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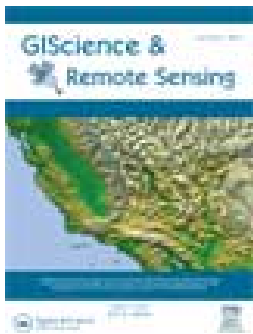
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# Peatland leaf-area index and biomass estimation with ultra-high resolution remote sensing

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## ABSTRACT

There is fine-scale spatial heterogeneity in key vegetation properties including leaf-area index (LAI) and biomass in treeless northern peatlands, and hyperspectral drone data with high spatial and spectral resolution could detect the spatial patterns with high accuracy. However, the advantage of hyperspectral drone data has not been tested in a multi-source remote sensing approach (i.e. inclusion of multiple different remote sensing datatypes); and overall, sub-meter-level leaf-area index (LAI) and biomass maps have largely been absent. We evaluated the detectability of LAI and biomass patterns at a northern boreal fen (Halssiaapa) in northern Finland with multi-temporal and multi-source remote sensing data and assessed the benefit of hyperspectral drone data. We measured vascular plant percentage cover and height as well as moss cover in 140 field plots and connected the structural information to measured aboveground vascular LAI and biomass and moss biomass with linear regressions. We predicted both total and plant functional type (PFT) specific LAI and biomass patterns with random forests regressions with predictors including RGB and hyperspectral drone (28 bands in a spectral range of 500–900 nm), aerial and satellite imagery as well as topography and vegetation height information derived from structure-from-motion drone photogrammetry and aerial lidar data. The modeling performance was between moderate and good for total LAI and biomass (mean explained variance between 49.8 and 66.5%) and variable for PFTs (0.3–61.6%). Hyperspectral data increased model performance in most of the regressions, usually relatively little, but in some of the regressions, the inclusion of hyperspectral data even decreased model performance (change in mean explained variance between –14.5 and 9.1%-points). The most important features in regressions included drone topography, vegetation height, hyperspectral and RGB features. The spatial patterns and landscape estimates of LAI and biomass were quite similar in regressions with or without hyperspectral data, in particular for moss and total biomass. The results suggest that the fine-scale spatial patterns of peatland LAI and biomass can be detected with multi-source remote sensing data, vegetation mapping should include both spectral and topographic predictors at sub-meter-level spatial resolution and that hyperspectral imagery gives only slight benefits.

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## Keywords

Biomass; hyperspectral imaging; leaf-area index; ultra-high spatial resolution; unmanned aerial systems (UAS)

## Introduction

Leaf-area index (LAI) and biomass are among the most important vegetation properties linked to biogeochemical processes (Wilson et al. 2007; van der Wal and Stien 2014). Green LAI quantifies the photosynthesizing leaf area per unit ground area and thus characterizes the interface between vegetation and atmosphere. It is a key parameter when the eddy covariance or chamber-based measurements of CO<sub>2</sub> and CH<sub>4</sub> are upscaled or modeled (Tuovinen et al. 2019; Wilson et al. 2007; Metzger et al. 2015). Plant biomass, in turn, refers to total mass of

plants; thus, it quantifies carbon stock in the vegetation and is closely linked to ecosystem productivity (Epstein et al. 2012; van der Wal and Stien 2014).

One way to characterize qualitative differences in vegetation is to assess LAI and biomass of different plant functional types (PFTs) (Juutinen et al. 2017; Yu et al. 2017). PFTs form a way to divide plant species into groups based on their growth forms and environmental responses (Duckworth, Kent, and Ramsay 2000; Ustin and Gamon 2010; Chapin et al. 1996). Users and their needs define the PFT grouping, but a common approach is to use groups such as evergreen and

deciduous shrubs, forbs, graminoids, and different types of mosses (Hugelius et al. 2011; Chapin et al. 1996; Räsänen et al. 2019). These PFTs differ in terms of how LAI and biomass are connected to plant percentage cover and height (Räsänen et al. 2019).

To capture the spatial patterns of LAI and biomass, remote sensing-based maps have been used for decades (Laidler and Treitz 2003; Walker et al. 2003). These maps can be PFT-specific (Greaves et al. 2016; Myers-Smith et al. 2015) or summed estimates of total LAI or biomass (Räsänen et al. 2019; Berner et al. 2018). Within treeless northern boreal or arctic environments, such as peatlands and tundra, the landscape is characterized by fine-scale spatial heterogeneity, controlled by variation in soil and bedrock characteristics, topography, and moisture. In particular, northern peatlands are usually flat but have notable heterogeneity in microtopography and vegetation evidenced by contrasting microforms, which often include wet and low-lying flarks, intermediate lawns and dry strings located higher than the other microforms (Laitinen et al. 2007). In these landscapes, ultra-high to very-high spatial resolution remote sensing datasets should be used (Virtanen and Ek 2014; Palace et al. 2018; Räsänen and Virtanen 2019), as the use of coarser-resolution maps can lead to erroneous landscape-level estimates of vegetation properties and associated biogeochemical processes (Räsänen and Virtanen 2019; Treat et al. 2018). Nevertheless, sub-meter-level LAI and biomass maps have rarely been developed for peatland landscapes, though such maps have been constructed in other environments such as temperate dryland (Cunliffe, Brazier, and Anderson 2016), shrub-dominated tundra (Cunliffe et al. 2020) and in particular in agricultural areas (Li et al. 2020; Yue et al. 2017; Poley and McDermid 2020).

In practice, ultra-high-resolution data can be gathered with unmanned aerial vehicles (i.e. drones). It also has been shown that there are multiple benefits when multi-source data is used, i.e. inclusion of multiple different datatypes collected from multiple different platforms, such as the combination of optical imagery and lidar data. In particular, with multi-source data, different aspects of the landscape, such as spectral, topographic, and vegetation height properties at multiple spatial resolutions can be captured. Thereby, the explanatory capacities are typically higher than when using a single data source (Räsänen and Virtanen 2019; Chen, Huang, and Xu 2017; Arroyo-Mora et al. 2017;

Luo et al. 2016; Sankey et al. 2018; Poley and McDermid 2020), and further boosts in model performance can be obtained with multi-temporal data (Halabisky, Babcock, and Moskal 2018; Gholizadeh et al. 2020; Poley and McDermid 2020).

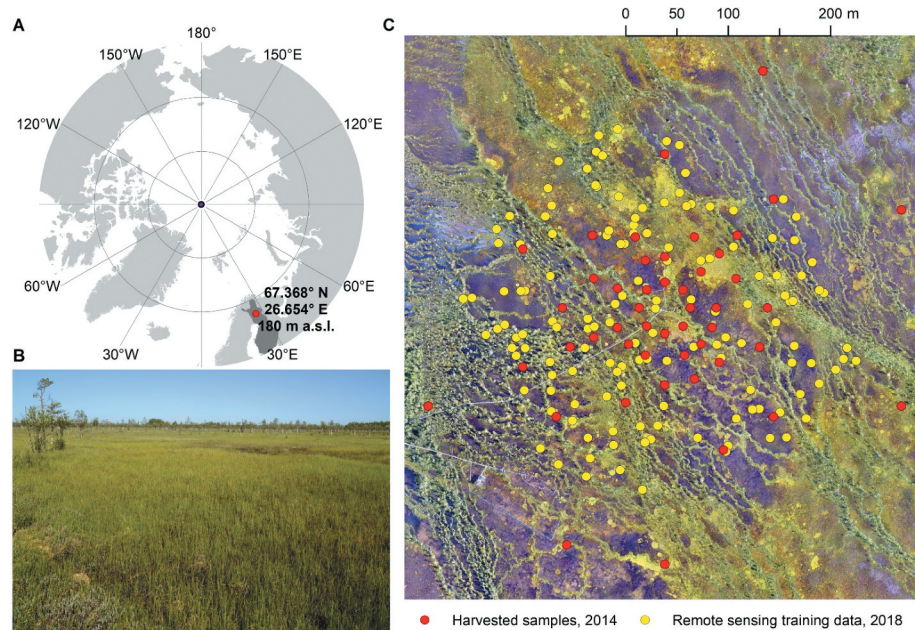
Although there are multiple different potential data sources, it has been discussed that some data such as hyperspectral imaging have multiple benefits as it can capture finer spectral signatures of vegetation than multispectral sensors (Chen, Chen, et al. 2017; Cole, McMorrow, and Evans 2013; Harris, Charnock, and Lucas 2015; Kalacska, Lalonde, and Moore 2015). In peatland landscapes, it has been shown that multiple vegetation properties can be tracked with hyperspectral data, for instance, floristic gradients (Harris, Charnock, and Lucas 2015), biotopes (Middleton et al. 2012), PFTs (Cole, McMorrow, and Evans 2013), and plant strategy types or plant traits (Kattenborn et al. 2017; Schmidtlein et al. 2012), including nitrogen and chlorophyll content (Kalacska, Lalonde, and Moore 2015). In spatially heterogeneous landscapes, the benefits of hyperspectral imaging are in particular evident with drone-based data, which can quantify the spectral properties of vegetation in high spatial and spectral resolutions (Aasen and Bolten 2018; Arroyo-Mora et al. 2019; Sankey et al. 2018; Aasen et al. 2015). However, relatively few studies have analyzed the advantages of hyperspectral drone data relative to other datasets in analyzing vegetation patterns in peatlands.

We address two major gaps in scientific knowledge, i.e. the lack of ultra-high-resolution LAI and biomass maps and the lack of testing the relative importance of hyperspectral drone data to map vegetation patterns in northern peatlands. We use multi-source and multi-temporal ultra-high-resolution remote sensing data to predict LAI and aboveground biomass patterns in a northern boreal fen landscape located in northern Finland. We ask the following two research questions. (1) How well can LAI and biomass patterns be detected with multi-source and multi-temporal ultra-high spatial resolution remote sensing data? (2) What is the benefit of hyperspectral drone data?

## Materials and methods

### Study area

Our study area is located in Halssiaapa fen in Sodankylä, northern Finland [Figure 1](#). The fen has



**Figure 1.** A) The location of study area in northern Finland, B) a photo from the study area, C) true-color drone image taken on 12 July 2016 with field inventory plots marked with dots.

fine-scale variation in vegetation, topography, wetness, and trophic status patterns (Dinsmore et al. 2017; Räsänen et al. 2019). The spatial heterogeneity can be characterized by three microforms: relatively low and dry strings (<0.5 m height) dominated by *Sphagnum* and evergreen and deciduous shrubs (e.g. *Andromeda polifolia* and *Betula nana*) and scattered birches (*Betula pubescens*) and pines (*Pinus sylvestris*); wetter flarks dominated by wet brown mosses (e.g. *Scorpidium* spp, *Warnstorfia exannulata*) and some graminoids (e.g. *Carex limosa* and *Eriophorum rufescens*); and intermediate lawns with continuous *Sphagnum* cover, forbs (e.g. *Menyanthes trifoliata*, *Comarum palustre*), graminoids (e.g. *Scheuchzeria palustris*) and some shrubs (e.g. *Vaccinium oxycoccos*). Trophic status varies from oligotrophic to eutrophic, and the transitions between different vegetation types and microforms are relatively gradual.

### Vegetation inventories

To collect ground and field layer LAI and aboveground biomass vegetation inventory data, we harvested samples from 48 square plots with 50 cm side length on July 17–19, 2014 (Figure 1). Species were classified into the following PFTs: evergreen dwarf shrubs, deciduous dwarf shrubs, forbs, graminoids, *Sphagnum* and other mosses. The PFT classification we used was a slight

modification of the one presented by Chapin et al. (1996). The plots were located along 250 m long transects to the cardinal, intercardinal, and secondary intercardinal directions from an eddy covariance flux tower located in the fen. We selected the plots so that all major land cover and vegetation types within the fen were included. We estimated the percentage cover of each PFT visually and measured the mean height of vascular plant PFTs using a ruler. We harvested aboveground parts of the plant species to measure LAI for vascular plants and biomass for vascular plants and mosses. We collected all the aboveground vascular plant material. For mosses, we collected a 5 cm x 5 cm sample of living green photosynthesizing part of the mosses. For measuring vascular LAI, we scanned the species or PFT-specific harvested samples with an A4 scanner and processed the scans with GIMP 2.8. software (The GIMP team, [www.gimp.org](http://www.gimp.org)). From each scanned image, we calculated the proportion of green pixels. For determining biomass, we oven dried the samples at 60°C for 24 hours (Klein and Bay 1994) and weighed them.

To collect training data for remote sensing-based regressions of LAI and biomass, we sampled 140 square plots with 50 cm side length using stratified random sampling on July 17–21, 2018 (Figure 1). Strata were four different land cover types from a multi-source remote sensing-based land cover map



(Mikola et al. manuscript in preparation). The plots were located at least 3 m apart from each other and the exact positions of the plots were collected with a Trimble R10 GPS device with  $\pm 5$  cm accuracy. At each plot, we evaluated the cover of vascular plant and moss species with visual interpretation and measured the mean height of vascular plant species with a ruler. To ensure the consistency in visual estimation of percentage cover, the last author inventoried part of the vegetation plots and trained the other surveyors during both field campaigns. Although there were some trees in the landscape, no trees were located in our field inventory plots; thus, we excluded trees (height  $>1.5$  m) in our analysis.

### Remote sensing datasets

The use of multi-source and multi-temporal remote sensing data has been documented to increase model performance in vegetation mapping, including biomass and LAI estimation (Halabisky, Babcock, and Moskal 2018; Chen, Huang, and Xu 2017; Sankey et al. 2018; Poley and McDermid 2020; Michez et al. 2018). Furthermore, it has been shown that optimal timing of remote sensing data depends on the studied ecosystem and target variable (Cole, McMorrow, and Evans 2014; Basnyat et al. 2004; Wang et al. 2019). Therefore, we used remote sensing data depicting the spectral, topography, and vegetation height properties captured from drone, aerial, and satellite platforms with 0.02-m to 3-m spatial resolution from multiple years and phenological stages (Table 1). In total, we calculated 359 features for each of the 140 field inventory plots. We calculated the mean value of each of the layers we used; in addition, we calculated the coefficient of variation for each hyperspectral band as well as a mean coefficient of variation over all bands (Wang et al. 2018).

We used hyperspectral drone images from two time points, one from the peak growing season (July 2019) and one from the beginning of senescence (late August 2019) Table 1, Figure 2. We acquired the imagery with the Senop Rikola (Senop Oy, Oulu, Finland) frame-based snapshot camera ( $1010 \times 1010$  pixels with  $36.5^\circ$  field of view) mounted on a DJI Matrice 600 hexacopter. We covered a 20 ha area with a 35–40 min flight time (11:10–11:45 AM on Jul 24, and 12:00–12:40 PM on Aug 28) at an altitude of 100 m and speed of  $15 \text{ km h}^{-1}$  ( $4.2 \text{ m s}^{-1}$ ) resulting

in ca. 1700 frames at 7 cm spatial resolution. The imagery was composed of 28 bands spanning the spectral range of 500–900 nm with an average bandwidth (full-width at half-maximum) of 8.9 nm (Table S1, Figure 2). We first converted the raw image data into radiance and applied a lens correction with the Rikola preprocessing software (Senop Oy, Oulu, Finland). As all the hyperspectral bands were acquired separately, the bands were not perfectly aligned and needed to be co-registered (Honkavaara et al. 2017; Jakob, Zimmermann, and Gloaguen 2017). We used the scale-invariant feature transform algorithm implemented in the MEPHySTo software for a practically perfect co-registration of the bands (Jakob, Zimmermann, and Gloaguen 2017). Then, we converted the images into reflectance and constructed mosaics with structure-from-motion (SfM) photogrammetry (Aasen et al. 2015) in Agisoft Metashape (Agisoft LLC, St. Petersburg, Russia). We conducted the reflectance conversion with empirical line calibration by utilizing three squared MosaicMill reflectance plates (MosaicMill Oy, Vantaa, Finland) with 0.5 m side length in 9%, 44%, and 75% reflectance. The plates were photographed during the flights. We georeferenced the image mosaics with 15 ground control points geolocated with a Trimble R10 GPS device with  $\pm 5$  cm accuracy. In addition to spectral bands, we calculated 80 different vegetation indices Table 2 with the *hsdar* package in R (Lehnert et al. 2019).

In addition to the hyperspectral imagery, we used true-color drone images from two time points, from a wetter (2016) and a drier year (2018) Table 1, Fig. S1. We acquired the images with a DJI Phantom 4 Pro ( $4000 \times 3000$  pixels with  $70^\circ/50^\circ$  field of view) and processed the data using SfM photogrammetry (Kalacska et al. 2017) in Pix4D (Pix4D S.A., Lausanne, Switzerland). On one hand, it has been shown that vegetation structure information is important in predicting biomass and LAI patterns (Poley and McDermid 2020; Cunliffe, Brazier, and Anderson 2016); on the other hand, peatland vegetation is largely controlled by micro-topographical variation (Räsänen and Virtanen 2019; Lehmann et al. 2016). Therefore, we produced 2-cm spatial resolution orthomosaics as well as a 9-cm spatial resolution digital terrain model (DTM) and vegetation height model (DTM subtracted from a digital surface model) Table 1. We georeferenced the 2018 image with 15 ground control points and co-registered the

**Table 1.** Remote sensing features used in leaf-area index and biomass regressions. Hyperspectral drone features are listed in Tables 2 and S1.

Dataset	Date	Producer	Spatial resolution	Number and list of layers
Hyperspectral drone images	24 July 2019; 28 August 2019	Authors	0.07 m	274: 28 bands, 80 indices, 29 coefficient of variation measures for both images
Drone image 1	12 July 2016	Authors	0.02 m	27: B, G, R, and 8 GLCM layers from all spectral bands
Drone image 2	11 July 2018	Authors	0.02 m	3: B, G, R
Drone digital elevation and terrain model	11 July 2018	Authors	0.09 m	6: Elevation, slope, TPIs (2 m and 5 m distance), TWI, VHM
Aerial image	19 August 2015	National Land Survey of Finland	0.5 m	4: B, G, R, NIR
Lidar data	19 August 2015	National Land Survey of Finland	0.5 points m <sup>-2</sup> (point cloud); 2 m (layers)	9: Elevation, slope, TPIs (5 m, 10 m, 20 m, 50 m, 100 m distances), TWI, VHM
PlanetScope images	May 16, Jun 2, Jun 17, Jul 3, Jul 10, Jul 19, Jul 27, Aug 8, Aug 26, Sep 9, Sep 18, 4 October 2018	Planet Labs Inc.	3 m	36: NDVI, NDWI, RGI from all images

Abbreviations: B, blue; G, green; GLCM, gray-level co-occurrence matrix; NDVI, normalized difference vegetation index; NDWI, normalized difference water index; NIR, near-infrared; R, red; RGI, red-green index; TPI, topographical position index; TWI, topographical wetness index; VHM, vegetation height model.

2016 image with the 2018 image using 20 control points.

In previous studies, it has been discussed that some of the LAI, biomass, and other vegetation patterns can be best detected with coarser resolution data (Poley and McDermid 2020; Räsänen et al. 2020). To detect coarser spatial resolution patterns of vegetation and topography, we included aerial lidar data and orthophotos. From the lidar data, we used a 2-m resolution DTM preprocessed by the National Land Survey and computed a vegetation height model by subtracting the DTM from a digital surface model in which we used all returns of the lidar point cloud. As vegetation spectral responses vary throughout the phenological cycle in peatlands (Cole, McMorrow, and Evans 2014), we used the surface reflectance product of 12 PlanetScope satellite images (Planet Team) taken between mid-May and early October 2018 Table 1.

Textural features have been shown to boost the accuracy in remote sensing-based mapping tasks (Mishra et al. 2018; Hall-Beyer 2017). Thus, for each spectral band of the 2016 drone image, we calculated eight gray-level co-occurrence matrix (GLCM) textural features (energy, entropy, correlation, inverse difference moment, inertia, cluster shade, cluster prominence and Haralick correlation) (Haralick, Dinstein, and Shanmugam 1973) with a moving window technique, eight quantization levels and neighborhood distance set to five in Orfeo Toolbox (Grizonnet et al. 2017). We computed the following topographical features from the drone

and lidar-based DTM: elevation, slope in degree, position index with different neighborhood radiuses (Guisan, Weiss, and Weiss 1999), and wetness index (Böhner and Selige 2006) in the SAGA GIS (Conrad et al. 2015). Instead of using the spectral bands as such for the PlanetScope images, we calculated the following band indices for each image: normalized difference vegetation index (NDVI) (Rouse et al. 1974), normalized difference water index (NDWI) (McFeeters 1996), and red-green index (RGI) (Coops et al. 2006) for each image Table 1.

### Data analysis

First, we established empirical relationships between (1) field-measured plant species cover and height, and (2) LAI and biomass for different PFTs using the harvested samples from 2014. We tested linear regressions with the following explanatory variable options: cover only and plant volume (cover × height), compared the performance of the two models and chose the final model based on root mean squared error (RMSE) value. We evaluated the final model performance with RMSE, normalized RMSE (nRMSE: i.e., RMSE divided by the difference between maximum observed value and minimum observed value), and adjusted coefficient of determination. We used the equations from linear regressions to predict PFT-specific LAI and biomass values for vegetation plots inventoried in 2018.

Second, for predicting LAI and biomass with remote sensing datasets, we conducted random

**Table 2.** Details of the high spectral resolution vegetation indices calculated for the hyperspectral data. In the equations, R refers to reflectance at a specific wavelength and D to first derivation of reflectance values at the wavelength.

Index	Equation	Reference
Boochs	$D_{703}$	Boochs et al. (1990)
Boochs2	$D_{720}$	Boochs et al. (1990)
CARI*	$R_{700} \times  a \times 670 + R_{670} + b  / R_{670} \times (a^2 + 1)^{0.5}$	Kim et al. (1994)
Carter2	$R_{695} / R_{760}$	Carter (1994)
Carter3	$R_{605} / R_{760}$	Carter (1994)
Carter4	$R_{710} / R_{760}$	Carter (1994)
Carter5	$R_{695} / R_{670}$	Carter (1994)
Carter6	$R_{550}$	Carter (1994)
CI	$R_{675} \times R_{690} / R_{683}^2$	Zarco-Tejada et al. (2003)
CI2	$R_{760} / R_{700} - 1$	Gitelson, Gritz, and Merzlyak (2003)
CIInt	$R_{735}$	Oppelt and Mauser (2004)
	$R_i$	
CRI1	$R_{515}^{-1} - R_{550}^{-1}$	Gitelson, Gritz, and Merzlyak (2003)
CRI2	$R_{515}^{-1} - R_{770}^{-1}$	Gitelson, Gritz, and Merzlyak (2003)
CRI3	$R_{515}^{-1} - R_{550}^{-1} \times R_{770}$	Gitelson, Gritz, and Merzlyak (2003)
CRI4	$R_{515}^{-1} - R_{700}^{-1} \times R_{770}$	Gitelson, Gritz, and Merzlyak (2003)
D1	$D_{730} / D_{706}$	Zarco-Tejada et al. (2003)
D2	$D_{705} / D_{722}$	Zarco-Tejada et al. (2003)
Datt	$(R_{850} - R_{710}) / (R_{850} - R_{680})$	Datt (1999)
Datt2	$R_{850} / R_{710}$	Datt (1999)
Datt3	$D_{754} / D_{704}$	Datt (1999)
Datt4	$R_{672} / (R_{550} \times R_{709})$	Datt (1998)
Datt5	$R_{672} / R_{550}$	Datt (1998)
Datt6	$R_{860} / (R_{550} \times R_{709})$	Datt (1998)
DD	$(R_{749} - R_{720}) - (R_{701} - R_{672})$	le Maire, François, and Dufrène (2004)
DDn	$2 \times (R_{710} - R_{660} - R_{760})$	le Maire et al. (2008)
DPI	$(D_{688} \times D_{710}) / D_{697}^2$	Zarco-Tejada et al. (2003)
DWSI4	$R_{550} / R_{680}$	Apan et al. (2004)
GDVI 2	$(R_{800}^2 - R_{680}^2) / (R_{800}^2 + R_{680}^2)$	Wu (2014)
GDVI 3	$(R_{800}^3 - R_{680}^3) / (R_{800}^3 + R_{680}^3)$	Wu (2014)
GDVI 4	$(R_{800}^4 - R_{680}^4) / (R_{800}^4 + R_{680}^4)$	Wu (2014)
GI	$R_{554} / R_{677}$	Smith et al. (1995)
Gitelson	$R_{700}^{-1}$	Gitelson, Buschmann, and Lichtenthaler (1999)
Gitelson2	$(R_{750} - R_{800} / R_{695} - R_{740}) - 1$	Gitelson, Gritz, and Merzlyak (2003)
GMI1	$R_{750} / R_{550}$	Gitelson, Gritz, and Merzlyak (2003)
GMI2	$R_{750} / R_{700}$	Gitelson, Gritz, and Merzlyak (2003)
Green NDVI	$(R_{800} - R_{550}) / (R_{800} + R_{550})$	Gitelson, Kaufman, and Merzlyak (1996)
Maccioni	$(R_{780} - R_{710}) / (R_{780} - R_{680})$	Maccioni, Agati, and Mazzinghi (2001)
MCARI	$((R_{700} - R_{670}) - 0.2 \times (R_{700} - R_{550})) \times (R_{700} / R_{670})$	Daughtry et al. (2000)
MCARI/OSAVI	MCARI/OSAVI	Daughtry et al. (2000)
MCARI2	$((R_{750} - R_{705}) - 0.2 \times (R_{750} - R_{550})) \times (R_{750} / R_{705})$	Wu et al. (2008)
MCARI2/OSAVI2	MCARI2/OSAVI2	Wu et al. (2008)
MPRI	$(R_{515} - R_{530}) / (R_{515} + R_{530})$	Hernández-Clemente, Navarro-Cerrillo, and Zarco-Tejada (2012)
MSAVI	$0.5 \times (2 \times R_{800} + 1 - ((2 \times R_{800} + 1)^2 - 8 \times (R_{800} - R_{670}))^{0.5})$	Qi et al. (1994)
mSR2	$(R_{750} / R_{705}) - 1 / (R_{750} / R_{705} + 1)^{0.5}$	Chen (1996)
MTCI	$(R_{754} - R_{709}) / (R_{709} - R_{681})$	Dash and Curran (2004)
MTVI	$1.2 \times (1.2 \times (R_{800} - R_{550}) - 2.5 \times (R_{670} - R_{550}))$	Haboudane et al. (2002)
NDVI	$(R_{800} - R_{680}) / (R_{800} + R_{680})$	Rouse et al. (1974)
NDVI2	$(R_{750} - R_{705}) / (R_{750} + R_{705})$	Gitelson and Merzlyak (1994)
NDVI3	$(R_{682} - R_{553}) / (R_{682} + R_{553})$	Gandia et al. (2004)
OSAVI	$(1 + 0.16) \times (R_{800} - R_{670}) / (R_{800} + R_{670} + 0.16)$	Rondeaux, Steven, and Baret (1996)
OSAVI2	$(1 + 0.16) \times (R_{750} - R_{705}) / (R_{750} + R_{705} + 0.16)$	Wu et al. (2008)
PARS	$R_{746} / R_{513}$	Chappelle, Kim, and McMurtrey (1992)
PRI	$(R_{531} - R_{570}) / (R_{531} + R_{570})$	Gamon, Peñuelas, and Field (1992)
PRI norm	$PRI / (RDVI \times R_{700} / R_{670})$	Zarco-Tejada et al. (2013)
PRI*CI2	$PRI \times CI2$	Garrity, Jan, and Vierling (2011)
PSSR	$R_{800} / R_{635}$	Blackburn (1998)
RDVI	$(R_{800} - R_{670}) / \sqrt{(R_{800} + R_{670})}$	Roujean and Breon (1995)
REP LE	Red-edge position through linear extrapolation	Cho and Skidmore (2006)
REP Li	$700 + 40 \times (((R_{670} + R_{780}) / 2 - R_{700}) / (R_{740} - R_{700}))$	Guyot and Baret (1988)
SAVI	$1.5 \times (R_{800} - R_{670}) / (R_{800} + R_{670} + 0.5)$	Huete (1988)
SPVI	$0.4 \times 3.7 \times (R_{800} - R_{670}) - 1.2 \times ((R_{530} - R_{670})^2)^{0.5}$	Vincini, Frazzi, and D'Alessio (2006)
SR	$R_{800} / R_{680}$	Jordan (1969)
SR1	$R_{750} / R_{700}$	Gitelson and Merzlyak (1997)
SR2	$R_{752} / R_{690}$	Gitelson and Merzlyak (1997)
SR3	$R_{750} / R_{550}$	Gitelson and Merzlyak (1997)
SR4	$R_{700} / R_{670}$	McMurtrey et al. (1994)
SR5	$R_{675} / R_{700}$	Chappelle, Kim, and McMurtrey (1992)
SR6	$R_{750} / R_{710}$	Zarco-Tejada and Miller (1999)
SR8	$R_{515} / R_{550}$	Hernández-Clemente et al. (2011)

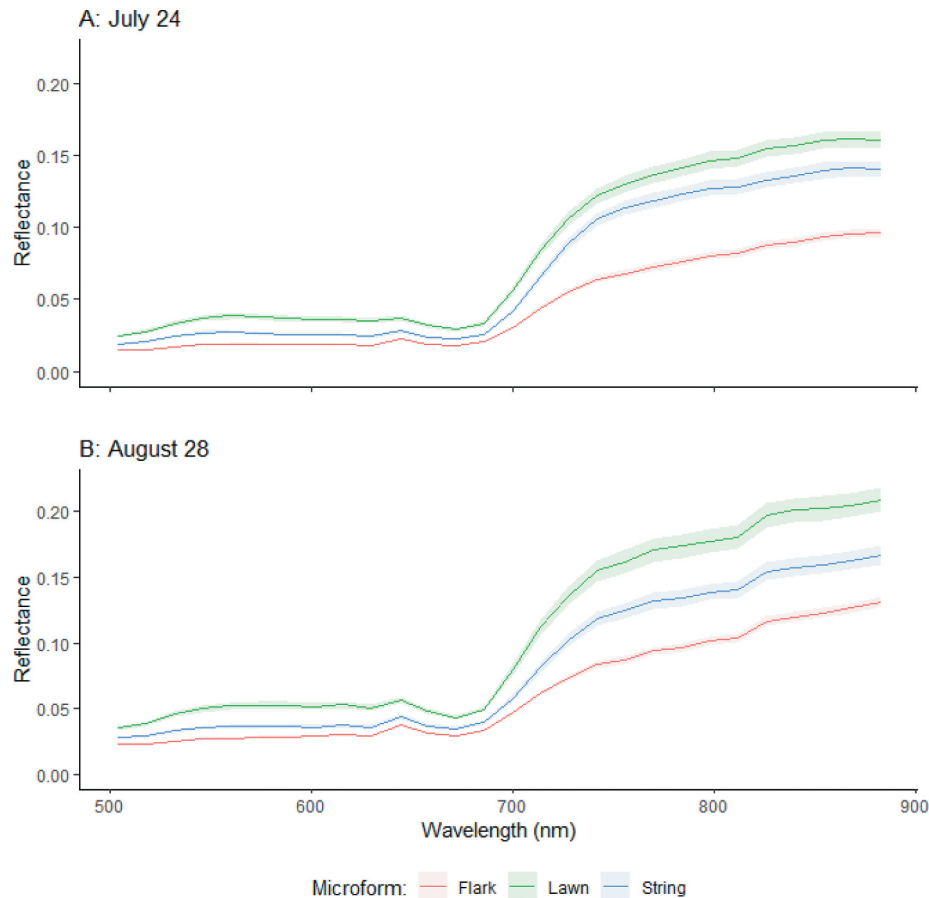
(Continued)



**Table 2.** (Continued).

Index	Equation	Reference
Sum Dr1	$\sum_{R_{626}}^{R_{795}}  D_{R_i} $	Elvidge and Chen (1995)
Sum Dr2	$\sum_{R_{680}}^{R_{780}} D_{R_i}$	Filella and Penuelas (1994)
TCARI	$3 \times ((R_{700} - R_{670}) - 0.2 \times (R_{700} - R_{550}) \times (R_{700} / R_{670}))$	Haboudane et al. (2002)
TCARI/OSAVI	$TCARI / OSAVI$	Haboudane et al. (2002)
TCARI2	$3 \times ((R_{750} - R_{705}) - 0.2 \times (R_{750} - R_{550}) \times (R_{750} / R_{705}))$	Wu (2014)
TCARI2/OSAVI2	$TCARI2 / OSAVI2$	Wu (2014)
TVI	$0.5 \times (120 \times (R_{750} - R_{550}) - 200 \times (R_{670} - R_{550}))$	Broge and Leblanc (2001)
Vogelmann	$R_{740} / R_{720}$	Vogelmann, Rock, and Moss (1993)
Vogelmann2	$(R_{734} - R_{747}) / (R_{715} + R_{726})$	Vogelmann, Rock, and Moss (1993)
Vogelmann3	$D_{715} / D_{705}$	Vogelmann, Rock, and Moss (1993)
Vogelmann4	$(R_{734} - R_{747}) / (R_{715} + R_{720})$	Vogelmann, Rock, and Moss (1993)

$$*a = (R_{700} - R_{550}) / 150; b = R_{550} - (a * R_{550})$$



**Figure 2.** Reflectance curves for the vegetation plots inventoried in 2018, based on the hyperspectral drone images taken on A) July 24 and B) 28 August 2019. The vegetation plots were classified into three different microforms. Lines indicate mean values and shaded region mean  $\pm$  standard error for the microforms.

forests (RF, Breiman 2001) regressions for the plots inventoried in 2018. RF has been used widely in biomass studies and it has often had the best goodness-of-fit (Poley and McDermid 2020); furthermore, it is insensitive to overfitting and can deal with a high

number of cross-correlated features typical to hyperspectral data (Belgiu and Dragut 2016). We conducted regressions for each PFT and separately for the following summed up estimates: LAI for vascular plants in total, and biomass for vascular plants in total, mosses

in total and all plant species in total. To test the added value of hyperspectral drone data in mapping LAI and biomass, we tested regressions with three different feature sets for each plant species grouping: (1) all features, (2) hyperspectral features only, and (3) all other features than hyperspectral features.

To remove redundant and non-important features from the models, we reduced the number of features with an RF wrapper feature selection algorithm called Boruta (Kursa and Rudnicki 2010) before the final RF regressions. We used the mean decrease accuracy feature importance measure and chose those features that were not rejected during 999 RF runs (i.e. their importance was not significantly lower than those of randomly shuffled shadow features). We evaluated the relative importance of different features with the feature importance scores from Boruta.

We conducted 100 RF regressions for each plant species grouping and feature set option. In each regression, we used 500 trees and set the number of tested features at each node to one-third of features in the model. For each set of 100 regressions, we calculated minimum, maximum, and mean values of (1) the amount of variance explained (pseudo  $R^2 = 1 - (\text{mean squared error})/(\text{variance}(\text{response}))$ ), (2) RMSE, and (3) nRMSE, with an out-of-bag evaluation in which approximately two-thirds of the data in each tree is used for training and the rest for evaluation (Breiman 2001). Earlier studies have shown that out-of-bag assessment gives an unbiased or even slightly conservative estimate of the model accuracy (Clark et al. 2010). Analyses were conducted in R (R Core Team 2018) using the packages randomForest (Liaw and Wiener 2002) and Boruta (Kursa and Rudnicki 2010).

We produced maps of LAI and biomass using a geographic object-based image analysis approach,

since such an approach eases the integration of different datasets and delineates the landscape into meaningful vegetation patches (Blaschke et al. 2014). We segmented the 2016 drone image with the multiresolution segmentation algorithm by Baatz and Schäpe (2000) implemented in TerraView (Câmara et al. 2008). Before the segmentation, we generalized the drone image to 5 cm spatial resolution using bilinear interpolation. We gave equal weight to color and shape information as well as to compactness and smoothness and set the similarity threshold to 0.12 resulting in segments with an average size of 6 m<sup>2</sup> (minimum size 3.3 m<sup>2</sup>). We then assigned LAI and biomass predictions of each RF model to each segment, estimated landscape-level LAI and biomass and compared the spatial pattern of LAI and biomass maps.

## Results and discussion

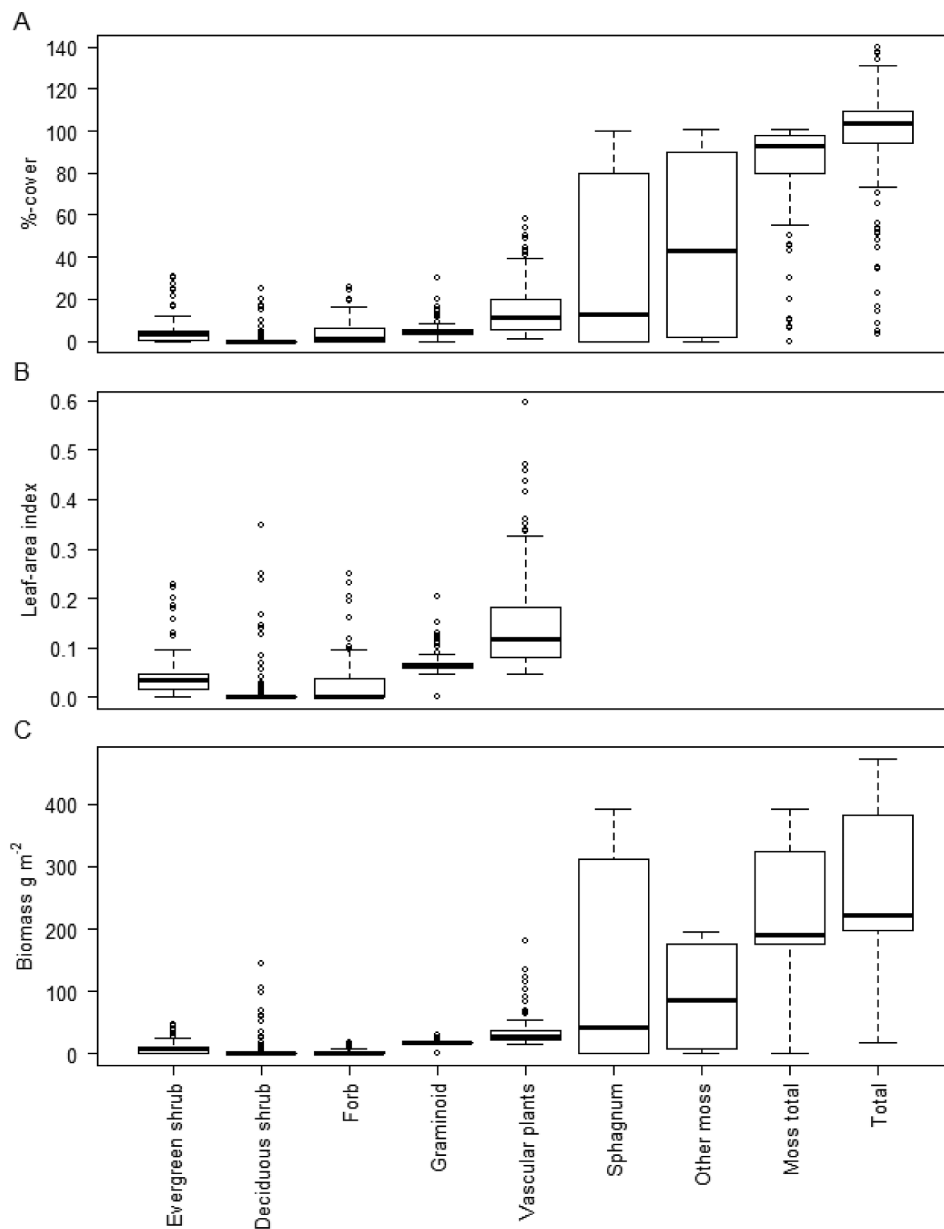
### Estimation of LAI and biomass from harvested samples

We could predict the PFT-specific LAI and biomass from samples harvested in 2014 with cover and height information with reasonably high accuracy (adjusted coefficient of determination between 0.50 and 0.98) Table 3. We applied these relationships to the plots inventoried in 2018 that were used as training data in remote sensing regressions. For the 2018 plots, mean LAI was 0.15 with the PFT-specific LAI estimate being the highest for graminoids Figure 3. Predicted biomass values were notably higher for mosses than for vascular plants, and the average value for total biomass was 260 g m<sup>-2</sup> Figure 3.

There was a four-year lag between our field inventories; hence, yearly variation might provide some

**Table 3.** Equations for calculating leaf-area index and biomass based on harvested samples ( $n = 48$ ), average and standard deviation values in the harvested data, and root mean squared error and adjusted coefficient of determination values.

Plant functional type	Leaf area index					Biomass				
	Equation	Mean $\pm$ sd (m <sup>2</sup> m <sup>-2</sup> )	RMSE (m <sup>2</sup> m <sup>-2</sup> )	nRMSE	R <sup>2</sup> <sub>adj</sub>	Equation	Mean $\pm$ sd (g m <sup>-2</sup> )	RMSE (g m <sup>-2</sup> )	nRMSE	R <sup>2</sup> <sub>adj</sub>
Evergreen shrub	LAI = 0.0138 + 0.0069*c	0.131 $\pm$ 0.148	0.052	0.09	0.87	BM = 3.51 + 1.35*c	26.4 $\pm$ 30.2	13.2	0.11	0.80
Deciduous shrub	LAI = -0.0012 + 0.0003*c*h	0.041 $\pm$ 0.127	0.018	0.02	0.98	BM = -0.63 + 0.12*c*h	17.1 $\pm$ 52.9	7.8	0.03	0.98
Forb	LAI = -0.0077 + 0.0010*c*h	0.078 $\pm$ 0.129	0.04	0.08	0.9	BM = -0.51 + 0.07*c*h	5.3 $\pm$ 8.9	3.2	0.07	0.86
Graminoid	LAI = 0.0420 + 0.0054*c	0.214 $\pm$ 0.099	0.064	0.15	0.56	BM = 15.40 + 0.45*c	30.0 $\pm$ 11.7	9.6	0.16	0.28
Sphagnum	n.a.	n.a.	n.a.	n.a.	n.a.	BM = -7.96 + 4.02*c	124.0 $\pm$ 182.8	83.0	0.11	0.79
Other moss	n.a.	n.a.	n.a.	n.a.	n.a.	BM = 4.36 + 1.89*c	88.7 $\pm$ 110.9	77.0	0.18	0.50
Total	n.a.	0.465 $\pm$ 0.325	n.a.	n.a.	n.a.	n.a.	291.5 $\pm$ 209.4	n.a.	n.a.	n.a.



**Figure 3.** Box and whisker plots for A) measured percentage cover, B) estimated leaf-area index, and C) estimated biomass values for each plant functional type and total in vegetation plots inventoried in 2018 ( $n = 140$ ).

uncertainties in our analysis. Nevertheless, during both field work years the relative abundance of different PFTs was similar (Table 3 and Figure 3) suggesting that yearly variation affects mostly the absolute biomass and LAI values and not their spatial patterns. Conversely, phenological stage may have a larger impact on the relative abundance of different PFTs (Juutinen et al. 2017; Cole, McMorro, and Evans 2014) but in our case both field inventories were conducted during peak growing season. Furthermore, field work data from 2014 were used only to link PFT cover and height to biomass and LAI. Such relationships do

not change rapidly in one study area and are also transferable and generalizable between study areas (Räsänen et al. 2019).

#### **Remote sensing of total and PFT-specific LAI and biomass**

In regressions between LAI and biomass and remote sensing data, the performance of the models varied markedly among different response variables (mean explained variance between  $-0.5\%$  and  $66.5\%$ , Table 4). Remote sensing regressions functioned well for

total vascular LAI and biomass and relatively well for total biomass but there was more variation in PFT-specific regressions [Tables 4 and 5](#).

Some PFTs could be predicted with relatively high accuracy (forbs, deciduous shrubs), and some with intermediate accuracy (evergreen shrubs). For graminoids, a low amount of variance was explained but RMSE was also low, while mosses had relatively high amount of explained variance and relatively high RMSE ([Tables 4 and 5](#)). On one hand, graminoids had low variance in LAI and biomass; i.e. there was a nearly contiguous graminoid cover in the landscape. On the other hand, mosses were abundant, could be observed from remote sensing data due to low vascular plant cover and had relatively high spatial variation of LAI and biomass [Figure 3](#).

Many studies have concentrated on mapping cover and biomass of shrubs, which are abundant in many northern treeless environments (Greaves et al. [2016](#); Myers-Smith et al. [2015](#)). In general, when a vegetation component is abundant and has relatively high variation in spatial patterns as well as distinct habitat and spectral and structural properties, it can be detected with remote sensing data more easily. There are notable differences between different northern treeless landscapes, including peatlands. In particular, when compared to our study area, some northern peatland landscapes have more (and some less) evident micro-topographical structure and higher (or lower) spatial variation vascular plant species abundance which is reflected in the ability to track the abundance and volume of different PFTs (Räsänen et al. [2020](#)). Therefore, one limitation of

our study was to concentrate only on one study area, and results could have been different in other sites.

### **The importance of hyperspectral and other remote sensing features**

There was clear PFT-specific variation how much model performance improved when hyperspectral data were included in ([Tables 4 and 5](#)). The benefit of using hyperspectral data was the most evident for evergreen shrubs; models with hyperspectral features only had slightly better performance than models with all features (0.2–0.4%-point higher mean explained variance), whereas the performance was lower for the models omitting hyperspectral features (5.6–9.1%-point decrease in mean explained variance). Also for vascular LAI (5.3%-point increase in mean explained variance) and biomass (2.8%-point increase), total biomass (2.1%-point increase), forbs, graminoids, other mosses, and mosses in total the inclusion of hyperspectral data increased model performance, but the models with hyperspectral features only had 6.1–23.6%-point lower mean explained variance. For *Sphagnum*, the models omitting hyperspectral features and with all features had approximately equal performance, but models with hyperspectral features only had 5.1%-point lower mean explained variance. For deciduous shrubs, models with hyperspectral data had a very low performance (mean explained variance 0.0% and 0.1% for LAI and biomass, respectively), but models with all features had a moderate performance (mean explained variance

**Table 4.** Explained variance (%) in random forests regressions for leaf-area index and biomass for different plant functional types and total. For each plant species grouping and feature set options, minimum, maximum and mean (in parentheses) explained variance over 100 random forests regressions are given.

Trait	Plant functional type	Feature set		
		All features	Hyperspectral only	Hyperspectral omitted
Leaf-area index	Evergreen shrub	31.9–36.9 (34.3)	32.1–37.4 (34.7)	25.3–31.6 (28.7)
	Deciduous shrub	42.2–49.7 (46.1)	–3.1–3.0 (0.0)	56.7–64.1 (60.6)
	Forb	49.9–54.0 (51.8)	26.7–32.1 (29.3)	47.7–52.4 (49.9)
	Graminoid	12.4–18.1 (15.3)	1.0–8.3 (4.7)	9.0–17.1 (14.2)
	Vascular total	65.1–68.4 (66.5)	57.7–61.2 (59.2)	59.5–62.7 (61.2)
Biomass	Evergreen shrub	34.7–39.7 (36.9)	35.2–39.5 (37.1)	24.7–31.2 (27.8)
	Deciduous shrub	45.0–52.7 (49.3)	–5.7–4.2 (0.1)	54.1–63.1 (58.2)
	Forb	50.7–55.1 (53.1)	32.6–38.8 (35.3)	46.6–51.1 (48.8)
	Graminoid	–0.1–0.6 (0.3)	–0.3–0.4 (0.1)	–1.0–0.2 (–0.5)
	Vascular total	60.8–65.9 (62.8)	35.9–42.2 (39.2)	57.5–62.3 (60.0)
	Sphagnum	60.4–62.9 (61.6)	55.3–58.2 (56.5)	59.7–62.9 (61.6)
	Other moss	55.3–58.1 (56.8)	46.1–48.8 (47.1)	54.5–57.5 (55.9)
	Moss total	47.3–51.4 (49.4)	40.2–43.4 (41.9)	44.8–48.5 (46.9)
	Total	47.2–51.1 (49.8)	41.3–45.6 (43.7)	47.3–50.7 (48.7)

**Table 5.** Root mean squared error (RMSE) and normalized RMSE (nRMSE) in random forests regressions for leaf-area index and biomass for different plant functional types and total. For each plant species grouping and feature set options, minimum, maximum and mean (in parentheses) RMSE and nRMSE over 100 random forests regressions are given.

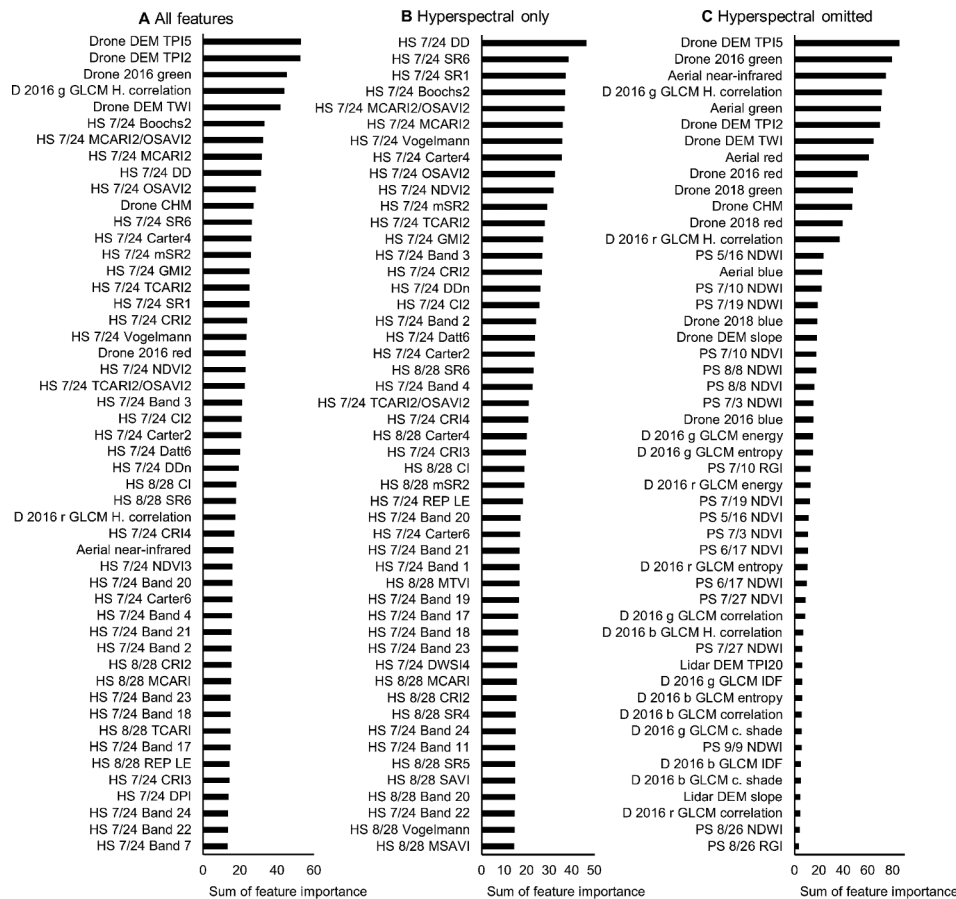
Trait	Plant functional type	Feature set					
		All features		Hyperspectral only		Excluding hyperspectral	
		RMSE	nRMSE	RMSE	nRMSE	RMSE	nRMSE
Leaf-area index	Evergreen shrub	0.035–0.037 (0.036)	0.15–0.16 (0.16)	0.035–0.036 (0.036)	0.15–0.16 (0.16)	0.037–0.038 (0.037)	0.16–0.17 (0.16)
	Deciduous shrub	0.034–0.035 (0.035)	0.10–0.10 (0.10)	0.047–0.049 (0.048)	0.13–0.14 (0.14)	0.029–0.031 (0.030)	0.08–0.09 (0.09)
	Forb	0.031–0.033 (0.032)	0.12–0.13 (0.13)	0.038–0.040 (0.039)	0.15–0.16 (0.16)	0.032–0.033 (0.033)	0.13–0.13 (0.13)
	Graminoid	0.022–0.023 (0.023)	0.11–0.11 (0.11)	0.023–0.024 (0.024)	0.11–0.12 (0.12)	0.022–0.023 (0.023)	0.11–0.11 (0.11)
	Vascular total	0.055–0.058 (0.056)	0.10–0.11 (0.10)	0.061–0.063 (0.062)	0.11–0.11 (0.11)	0.060–0.062 (0.061)	0.11–0.11 (0.11)
Biomass	Evergreen shrub	6.9–7.2 (7.1)	0.15–0.16 (0.16)	6.9–7.2 (7.1)	0.15–0.16 (0.16)	7.4–7.7 (7.6)	0.16–0.17 (0.17)
	Deciduous shrub	13.7–14.7 (14.1)	0.09–0.10 (0.10)	19.4–20.4 (20.0)	0.13–0.14 (0.14)	12.1–13.4 (12.8)	0.08–0.09 (0.09)
	Forb	2.1–2.2 (2.1)	0.13–0.13 (0.13)	2.4–2.6 (2.5)	0.14–0.15 (0.15)	2.2–2.3 (2.2)	0.13–0.14 (0.13)
	Graminoid	2.4–2.5 (2.5)	0.08–0.09 (0.09)	2.5–2.5 (2.5)	0.09–0.09 (0.09)	2.5–2.6 (2.6)	0.09–0.09 (0.09)
	Vascular total	13.8–14.8 (14.4)	0.08–0.09 (0.09)	17.9–18.9 (18.4)	0.11–0.12 (0.11)	14.5–15.4 (14.9)	0.09–0.09 (0.09)
	Sphagnum	95.7–98.9 (97.3)	0.24–0.25 (0.25)	101.6–105.0 (103.6)	0.26–0.27 (0.26)	95.7–99.8 (97.3)	0.24–0.25 (0.25)
	Other moss	49.9–51.5 (50.6)	0.25–0.26 (0.26)	55.1–56.6 (56.0)	0.28–0.29 (0.29)	50.2–52.0 (51.1)	0.26–0.27 (0.26)
	Moss total	75.5–78.6 (77.0)	0.19–0.20 (0.20)	81.5–83.7 (82.5)	0.21–0.21 (0.21)	77.7–80.4 (78.9)	0.20–0.20 (0.20)
	Total	78.8–81.9 (79.9)	0.17–0.18 (0.18)	83.1–86.3 (84.6)	0.18–0.19 (0.19)	79.2–81.7 (80.7)	0.17–0.18 (0.18)

46.1% and 49.3% for LAI and biomass, respectively), and models omitting hyperspectral features even better performance (58.2% and 60.6% for LAI and biomass, respectively) [Table 4](#).

In the models with all features, peak growing season vegetation indices calculated from hyperspectral data, drone topography, wetter year drone green and red band, GLCM Haralick correlation and drone vegetation height were among the most important ones [Figure 4a](#). In the models with hyperspectral features only, peak growing season features were typically more important than late summer features but also late summer features were deemed important in many regressions [Figure 4b](#). When we omitted hyperspectral data from the regressions, drone spectral, textural, topographic, and vegetation height features were among the most important but also aerial imagery features, in particular the near-infrared band, and early and midsummer PlanetScope features were among the most important ones [Figure 4c](#). Hyperspectral coefficient of variation, lidar, and late summer PlanetScope features had low importance and were often rejected by Boruta.

Our hyperspectral data included only 28 spectral bands in a relatively narrow spectral range (500–900 nm). If the data would have included more bands with narrower bandwidth and wider spectral range in total, the results could have been different and the importance of hyperspectral data more evident. Nonetheless, in a previous study by Harris, Charnock, and Lucas (2015) which utilized airborne hyperspectral data for mapping peatland vegetation gradients, it was found that relatively few bands (12 or 16) were selected for final regression models. In another study, Turner et al. (2019) assessed optimal spatial and spectral resolution for mapping health of Antarctic moss vegetation with different drone-based multispectral and hyperspectral sensors and concluded that 25 VNIR bands at 8 cm spatial resolution was an ideal band constellation. These findings suggest that hundreds of bands are not necessarily needed when mapping vegetation patterns in peatlands and similar landscapes. However, if the goal is to identify individual plant species instead of PFTs, better spectral resolution might aid more as some





**Figure 4.** Sum of Boruta feature importance scores for the 50 most important features over all regressions with A) all features, B) hyperspectral features only, and C) hyperspectral features omitted. In the figure, HS refers to hyperspectral, other abbreviations are listed in Tables 1, 2 and S1.

species have their own specific spectral signatures (Falcioni et al. 2020).

Our results suggest that some of the differences in vegetation cannot be delineated with spectral data; instead, also structural and topographic features are needed to obtain the highest accuracies Tables 4 and 5. Similar results have been obtained also previously in studies utilizing hyperspectral data (Arroyo-Mora et al. 2017; Luo et al. 2016; Sankey et al. 2018; Yue et al. 2017; Li et al. 2020). This is evident in particular with deciduous shrubs, which could not be mapped with hyperspectral data only; i.e. the models had very low explanatory capacities Table 4 and predicted values relatively high variance Table 6. Instead, for this PFT, the most important remote sensing features included topographic and vegetation height features derived from SfM drone data. This suggests that deciduous shrub abundance follows micro-topographic variation within the fen landscape, with the deciduous shrubs being most abundant in strings.

Therefore, our results highlight that both topographic and spectral features should be used when mapping vegetation in patterned fen landscapes. Nevertheless, not all vegetation follows the micro-topography and there is also a notable variation of vegetation (and thus spectral signatures) within the different micro-forms; part of which can be tracked with high-resolution spectral data. This is evident in particular with evergreen shrubs, for which the highest explanatory capacities were obtained with hyperspectral features only Table 4.

According to our results, a majority of the most important features were calculated from ultra-high resolution datasets (spatial resolution <10 cm), and the less fine-resolution data with a spatial resolution of 0.5–3.0 m were not as important Figure 4. Similar patterns could be observed both for spectral and structural/topographic features: data from aerial and satellite platforms were of less importance than drone data. Therefore, our results support previous findings

**Table 6.** Landscape-level mean ( $\pm$  standard deviation) leaf-area index and biomass for plant functional type-specific and total estimates based on maps produced with different feature sets.

Trait	Plant functional type	Feature set		
		All features	Hyperspectral only	Hyperspectral omitted
Leaf-area index	Deciduous shrub	0.012 $\pm$ 0.013	0.014 $\pm$ 0.025	0.012 $\pm$ 0.013
	Evergreen shrub	0.038 $\pm$ 0.018	0.048 $\pm$ 0.024	0.036 $\pm$ 0.02
	Forb	0.034 $\pm$ 0.031	0.033 $\pm$ 0.024	0.026 $\pm$ 0.032
	Graminoid	0.069 $\pm$ 0.007	0.070 $\pm$ 0.012	0.070 $\pm$ 0.008
	Vascular total	0.166 $\pm$ 0.062	0.187 $\pm$ 0.071	0.149 $\pm$ 0.053
Biomass (g m <sup>-2</sup> )	Deciduous shrub	5.3 $\pm$ 5.9	5.7 $\pm$ 9.5	4.7 $\pm$ 5.8
	Evergreen shrub	8.2 $\pm$ 4.1	8.1 $\pm$ 4.1	8.1 $\pm$ 4.2
	Forb	2.4 $\pm$ 2.2	2.4 $\pm$ 1.8	1.8 $\pm$ 2.2
	Graminoid	17.7 $\pm$ 0.6	17.7 $\pm$ 0.7	17.1 $\pm$ 0.6
	Vascular total	35.1 $\pm$ 11.6	39.5 $\pm$ 16.4	33.1 $\pm$ 10
	Sphagnum	104.5 $\pm$ 109.4	100.2 $\pm$ 104.4	112.9 $\pm$ 112.1
	Other moss	99.2 $\pm$ 55.3	100.9 $\pm$ 53.4	97.1 $\pm$ 52.8
	Moss total	198.9 $\pm$ 74.8	200.4 $\pm$ 69.6	211.8 $\pm$ 64.7
	Total	236.3 $\pm$ 78.6	235.5 $\pm$ 76.9	247.9 $\pm$ 71.3

that drone-derived DTMs can reveal fine-scale variation in micro-topography which controls vegetation patterns and ecosystem functioning in flat peatland landscapes (Harris and Baird 2019; Moore et al. 2019), and drone imagery can be used to track peatland vegetation (Palace et al. 2018; Beyer et al. 2019). Yet, the importance of ultra-high-resolution data is contradicted by our earlier results indicating that as high land cover mapping accuracies can be achieved with 50 cm spatial resolution data than with <10 cm spatial resolution data in a relatively similar type of peatland landscape (Räsänen and Virtanen 2019). However, in that study, only RGB drone imagery instead of hyperspectral data were used and land cover instead of biomass and leaf-area index was mapped. Although our study highlighted the importance of ultra-high-resolution datasets, also coarser-resolution features were among the most important in some regressions Figure 4, in particular in regressions omitting hyperspectral data. In these regressions, aerial imagery and PlanetScope provided spectral information with an infrared band that could not be obtained with RGB drone data.

We utilized only one ultra-high resolution vegetation structural feature (mean vegetation height based on SfM photogrammetry), which was important in particular for deciduous shrub LAI and biomass prediction. However, its value was more limited for other regressions. In the visual interpretation of the vegetation height model, it could be seen that the height of forbs and graminoids was not accurately predicted, and higher values for vegetation height were located in strings with higher shrub cover; moreover, the location of trees in strings was quite well predicted.

In earlier studies, SfM-based vegetation height models have been used, for example, in predicting biomass patterns in temperate arid environments with some trees, tall shrubs and grass tussocks (Cunliffe, Brazier, and Anderson 2016), biomass patterns in shrub-dominated tundra (Cunliffe et al. 2020), and in estimating vegetation height of shrubs in low-arctic tundra (Fraser, Ian Olthof, and Schmitt 2016). In all of these studies, the spatial resolution has been  $\leq 1$  cm compared to 9 cm in our study. A finer spatial resolution could thus help in achieving a more accurate prediction of height and volume of different PFTs, and from a finer resolution data, more features than only mean height could have been quantified. Even better results could be achieved with terrestrial or ultra-high resolution aerial/drone-based lidar (Greaves et al. 2016, 2017; Sankey et al. 2018), which has outperformed SfM in prediction accuracy in forested areas due to the capability to map terrain elevation below dense canopies (Wallace et al. 2016). Therefore, future research avenues could include mapping of ultra-high resolution of vegetation structure for different PFTs, and comparison of different methods such as SfM and lidar in open treeless environments. However, in peatlands where a majority of total biomass is composed of mosses, structural features should be coupled with spectral features.

Only a part of the remote sensing data was from the same year when the field inventory training data for remote sensing was collected (i.e. 2018, Table 1), which could potentially lead to errors in LAI and biomass prediction. Despite this fact, RGB drone features from the field work year of 2018 had lower importance in the regressions than RGB drone features

from the wetter year of 2016 [Figure 4](#), when vegetation was more abundant and there was higher contrast in the image. As both images were taken during the peak growing season, it can be reasoned that remote sensing data may not need to be from the same year if it is taken during similar phenological stage. In previous research in peatland landscapes, it has been shown that during peak growing season, vegetation volume is at its highest and plant functional types are spectrally separable from each other (Cole, McMorrow, and Evans 2014) and that spatial patterns of biomass and LAI vary throughout the phenological cycle (Juutinen et al. 2017). According to our feature importance results in [Figure 4](#), peak growing season hyperspectral and PlanetScope features were more important than non-peak season features. However, also non-peak season features and RGB drone features from two different years (2016 and 2018) were important in the different regressions [Figure 4](#). This indicates that multitemporal data can alleviate problems related to temporal mismatches and increase model performance (Räsänen et al. 2020; Halabisky, Babcock, and Moskal 2018; Gholizadeh et al. 2020; Poley and McDermid 2020). Nevertheless, there is some uncertainty when comparing the relative benefit of datasets with divergent spectral and spatial resolutions as the data were acquired at different times; in an ideal situation, all datasets would have been collected simultaneously. Multi-temporal datasets could also be integrated with temporal models utilizing environmental parameters such as maximum ecosystem photosynthesis or climatic data but the use of such models would require also multi-temporal vegetation inventories (Juutinen et al. 2017).

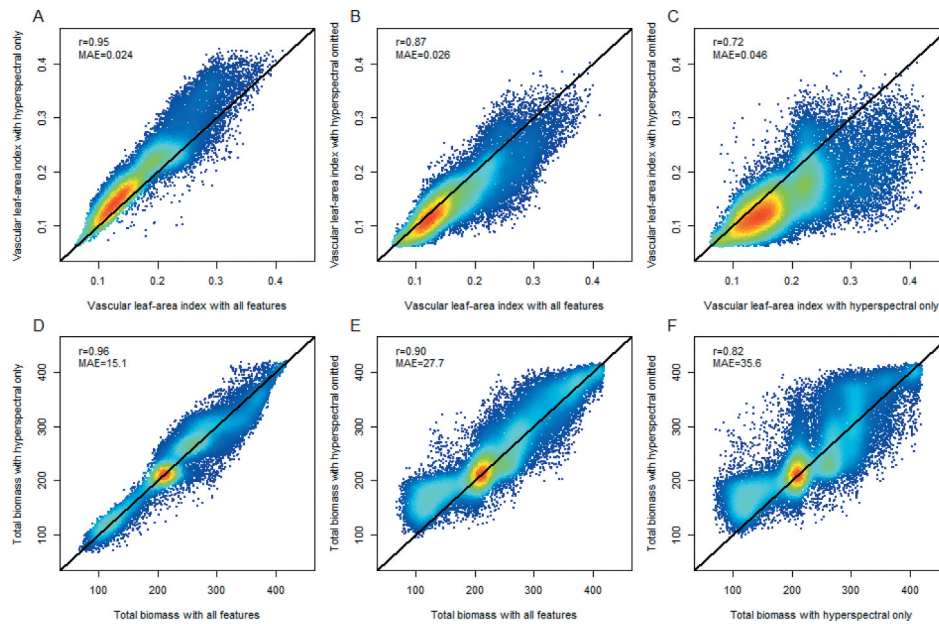
When looking at the reflectance spectra of the hyperspectral images [Figure 2](#), reflectance at all wavelengths was higher in the late growing season than during peak season which might indicate that conversion to absolute reflectance values was imperfect with the reflectance plates, which might affect results of the benefit of hyperspectral data. However, the reflectance differences between the images were higher in the visible spectrum, and therefore, consistent with earlier literature, spectral indices such as NDVI, MERIS terrestrial chlorophyll index (MTCI, Dash and Curran 2004) and red-edge position (Cho and Skidmore 2006) dropped between peak season and late season showing the start of senescence (Cole, McMorrow, and Evans 2014; Harris and Dash 2011). The drop was more evident in strings that have higher

vascular plant cover than moss-dominated lawns and flarks indicating that leaf senescence of deciduous vegetation and decrease in chlorophyll affected vascular plants more than mosses. This finding is backed by previous research in which it has been found that phenology derived changes in reflectance are higher in shrubs and other vascular plants than in mosses (Cole, McMorrow, and Evans 2014).

### ***Landscape-level estimates of LAI and biomass***

Overall, LAI and biomass estimates with remote sensing regressions with three different feature sets were typically quite close to each other, yet there was some variation between patches and landscape in total [Table 6](#), [Figures 5](#) and [6](#). The differences in estimates were relatively small between the three different feature set options (all data, hyperspectral only, hyperspectral omitted) when we predicted total or moss biomass [Table 6](#), [Figure 5\(d-f\)](#), [Figures 6\(f-h\)](#). There was only some overestimation in regressions omitting hyperspectral data. In *Sphagnum* biomass regressions, hyperspectral regressions had slight underestimation and regressions without hyperspectral features overestimation [Table 6](#). As also the explanatory capacities were relatively high in moss and total biomass regressions, it can be suggested that all three feature set options can be used for operational purposes. For vascular plants, the differences in plot and landscape-level estimates were higher [Table 6](#), [Figures 5\(a-c\)](#), [Figures 6\(a-c\)](#). When compared to models with all features, regressions with hyperspectral data only overestimated in particular deciduous shrub and total LAI and biomass as well as evergreen shrub LAI, whereas regressions omitting hyperspectral data underestimated forb and total LAI and biomass as well as deciduous shrub biomass [Table 6](#).

Vascular LAI patterns followed the micro-topographical variation in the fen landscape with the highest values being in the strings [Figures 6\(a-c, e\)](#), whereas for total biomass, the highest values were found in those lawn and string patches that had continuous moss cover [Figures 6\(f-h\)](#). When our ultra-high resolution biomass maps were compared with biomass maps produced with 2 m spatial resolution satellite imagery in the same site (Räsänen et al. 2019), the string-flark pattern was better distinguished on the finer-resolution maps, further suggesting the benefits of using ultra-high spatial resolution data.



**Figure 5.** Pairwise map comparisons for vascular leaf-area index (A–C) and total biomass (D–F) for regressions with different feature sets. Each dot in the plots represents LAI or biomass prediction for one segment. Dot density is showed with heat map colors so that red indicates higher density and blue lower density. In the figure,  $r$  refers to Pearson correlation coefficient and MAE to mean absolute error between the maps.

Our overall estimates of vascular LAI and biomass Figures 3, 5 and 6 were lower compared to previous coarser resolution estimates in the study area (Räsänen et al. 2019) and other estimates in similar peatland landscapes (Wilson et al. 2007; Laine et al. 2019). This is related to the dry and warm year in 2018 (Fig. S1) during which the cover and volume of vascular plants were low and the training data for remote sensing analyses were collected. However, as already discussed in Section “Estimation of LAI and biomass from harvested samples” the yearly variation affected the total abundance and volume of different PFTs but not their relative abundance nor spatial patterns. Thus, we judge that the absolute LAI and biomass values could have been better predicted remote sensing data concurrent with field work but the spatial patterns can be well predicted with asynchronous data.

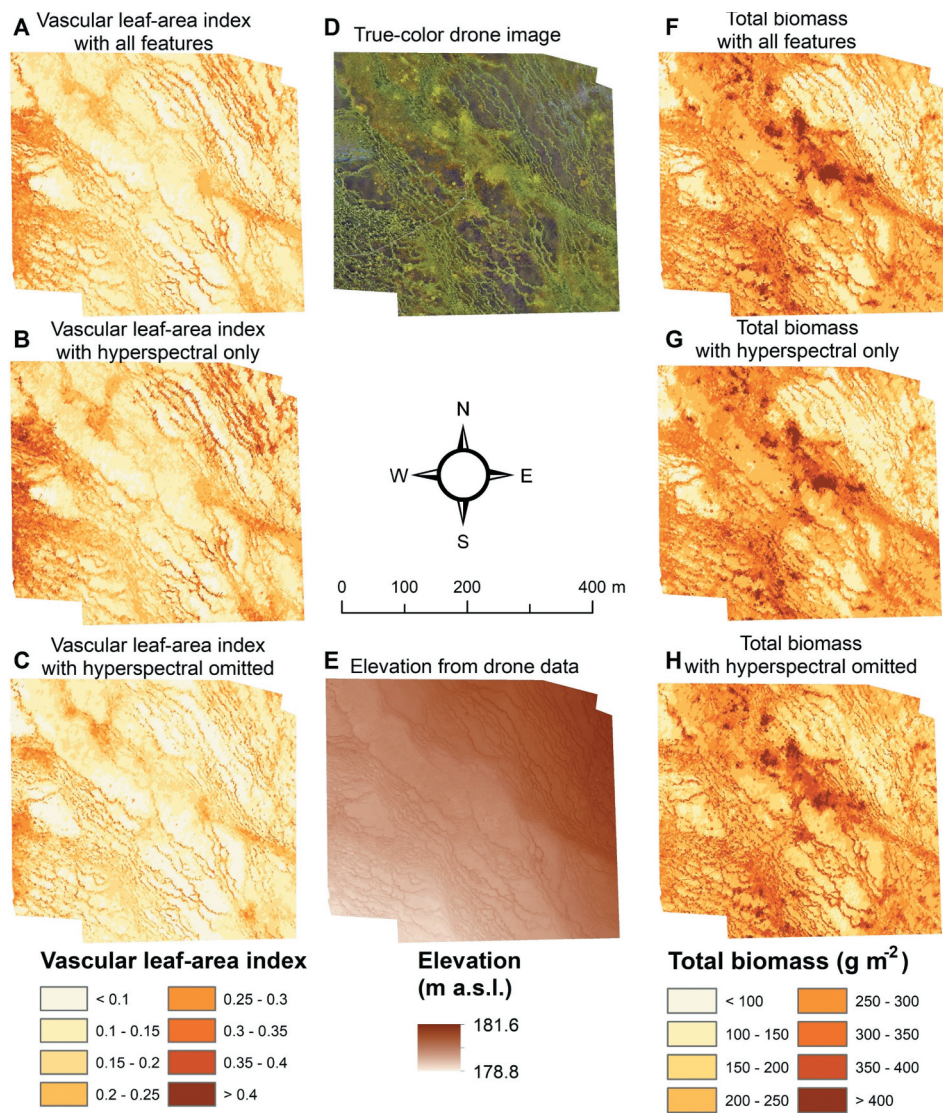
Our analysis excluded trees as we did not have suitable training data for predicting tree LAI and biomass. Few trees found in the study area are growing only in strings and are a minor component of vegetation; moreover, as there were no trees in our field plots, they had no impact on our model parameters. If the goal would be the prediction of tree LAI and biomass, it should be pursued separately from other PFTs in particular in these kinds of landscapes with few trees, and in

the prediction, ultra-high-resolution vegetation height model should be utilized (Cunliffe, Brazier, and Anderson 2016; Cunliffe et al. 2020; Wallace et al. 2016).

## Conclusions

We evaluated how well-peatland LAI and biomass patterns can be detected with multi-source and multi-temporal ultra-high-resolution remote sensing data and examined the benefit of hyperspectral drone data. The remote sensing-based regression models for total vascular LAI and biomass had a relatively high explanatory capacity, and also the model for total biomass functioned moderately well, suggesting good detectability of LAI and biomass patterns with ultra-high resolution remote sensing in spatially heterogeneous landscapes such as northern peatlands. There was more variation in PFT-specific models. As explanatory capacities were higher for total LAI and biomass than for PFT-specific estimates, our results suggest that maps of total LAI and biomass should not be summed maps of PFT-specific estimates but the response variable in the regressions and maps should be the total LAI or biomass. These maps could then be complemented with maps for such





**Figure 6.** Maps of vascular leaf-area index (A–C) and total biomass (F–H) with different feature sets used as the input data. Additionally, a true-color drone image from 2016 (D) and elevation based on drone data (E) are included.

PFTs that can be mapped with relatively high accuracy.

Our results showed mixed results on whether hyperspectral drone data are important in tracking LAI and biomass patterns. According to feature importance scores, hyperspectral data were extremely important; yet, when looking at explanatory capacities of different models, the inclusion of hyperspectral data boosted performance in most models but usually rather modestly. Additionally, landscape-level LAI and biomass estimates regressions with three different feature sets (all features, hyperspectral features, hyperspectral omitted) were typically quite close to each other. These results suggest that hyperspectral data give only slight benefits when included

in a multi-source remote sensing analysis of LAI and aboveground biomass patterns. In future studies, the relative benefit of different multispectral and hyperspectral sensors with various band constellations and spectral ranges could be tested in different landscapes.

Our results showed that multiple different remote sensing datasets are important in LAI and biomass estimation; hence, we suggest that (hyper)spectral datasets should be combined with topographic and structural data at sub-meter-level spatial resolution when mapping vegetation patterns in spatially heterogeneous landscapes such as northern peatlands. We also found that peak growing season features were more important than non-peak season features



but more research should be conducted for evaluating the optimal timing for remote sensing data acquisition.

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## Data availability

The data that support the findings of this study are available from the corresponding author, AR, upon reasonable request.

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